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PATENT SPECIFICATION

M Robin

Inventors: ROBERT PALMER RUH and RALPH ANDERSON DAVIS



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COMPLETE SPECIFICATION

Improved Chromium Fluoride Catalyst and Catalytic Process

We, THE DOW CHEMICAL COMPANY, 2 Corporation organized and existing under the Laws of the State of Delaware, United States of America, of Midland, County of Midland, State of Michigan, United States of America. do hereby declare the invention, for which we pray that a Patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the 10 following statement: —

This invention relates to an improved fluorination catalyst, and to a process for fluorinating halohydrocarbons to highly fluorinated products with the aid of this

15 catalyst.

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Heretofore, it has been known that chromium fluoride is useful in promoting the vapour-phase fluorination reaction of hydrogen fluoride with certain haloalkanes at elevated temperatures above 350°C. In British Patent No. 468,447, Leicester discloses that chromic fluoride supported on carbon in a massive or granular form is a suitable catalyst in such a process. This catalyst is shown to be effective 25 at temperatures between 450°C, and 550°C. in catalyzing the reaction of carbon tetrachloride with hydrogen fluoride to form CCl₃F and CCl₂F₂ as the chief reaction products. Although Leicester obtained only 8 per 30 cent CCIF, in carrying out this reaction at 550°C. he indicates that CF4 may be formed at this same temperature if a large proportion of hydrogen fluoride to carbon tetrachloride is employed.

This supposition, however, has recently been controverted by United States Patent No. 2,458,551 granted to Benning et al who found that CrF, supported on activated carbon, or pellets of CTF, per se, will not effectively 40 catalyze the reaction of HF with CCl, to form CF, unless a relatively high temperature in the range of from 700°C, to about 1000°C, is employed. In fact, I maing et al show that at

[Price 3s. 6d.]

temperatures below about 700°C., only insignificantly small amounts of CF, are produced even if HF is present in an amount up to 300 per cent in excess of that theoretically required to prepare CF₁. For example, when HF and CCl₂F₂ in a mole ratio of 3.9:1 were passed over a CrF, catalyst at 674°C., only 0.4 per cent conversion to CF, was obtained.

While the catalysts set forth in the above

Patent Specifications have been recommended

broadly for the reaction of chloromethanes and hydrogen fluoride, they are effective mainly in

inducing the formation of fluorinated products containing a low degree of fluorine substitution. These catalysts are even less effective in the reaction of hydrogen fluoride with bromoalkanes than with chloroalkanes due to excessive decomposition. For this reason, the production of highly fluorinated bromoalkanes,

such as bromotrifluoromethane, by vapourphase reactions with hydrogen fluoride has not been commercially practicable. In fact, it is only recently that CF, Br has been prepared

by any method.

The present invention is an advance over these known practices. It is based on the discovery that hydrated chromium fluoride may be activated with oxygen as hereinafter particularly described, and that the material so activated is very effective in catalyzing the vapour-phase fluorination reaction of haloalkanes and hydrogen fluoride. In fact, the catalysis of the invention, believed to contain a novel basic chromium fluoride, are more active than CrF,, or any of the catalysis known in the literature. They are also more effective in directing the course of the vapour-phase fluorination to greater conversions and vields of more highly fluorinated products, and at much lower temperatures, than has heretofore been achieved. For example, at a temperature as low as 150°C., CCI, and HF are reacted preponderantly to CCl₂F₂, while at a slightly

higher temperature e.g. 250°C., CCl₄ is fluorinated predominantly to CF₃Cl and CF₄. It has also been discovered that bromoalkanes react readily with hydrogen fluoride over these new catalysts to produce highly fluorinated bromoalkanes, i.e. CBr₄ is fluorinated to CF₃Br.

The compounds which may be fluorinated using catalysts according to the invention are partially or completely halogenated aliphatic hydrocarbons containing no more than four carbon atoms, no lodine, and at least one halogen other than fluorine. More specifically, the aliphatic halohydrocarbon reactants have from one to four carbon atoms including at least one carbon atom which is attached to a minimum of two halogens of atomic number not greater than 35, viz., fluorine, chlorine, or bromine, at least one of said halogens being of atomic 20 number from 17 to 35 inclusive, viz., chlorine or bromine. The aliphatic halohydrocarbon reactant is, therefore, a chloro-, bromo-, or chlerobromo- substituted product of a hydrocarbon (or fluorohydrocarbon). When the halohydrocarbon reactant centains hydrogen atoms and two or more carbon atoms in the molecule, from the standpoint of thermal stability it is desirable that all of the halogen atoms be attached to a single carbon atom. In general, 30 saturated halohydrocarbons are employed as reactants in the present process, preferably having one to two carbon atoms per molecule or haloalkanes having from one to two carbon atoms and from three to four chlorine or bromine atoms attached to a single carbon atom, e.g. carbon tetrachloride, chloroform, or 1,1,1-trichloroethane or the bromine analogues thereof. Other halomethanes which are fluorinated by the process of the invention inciude methylene chloride, methylene bromide, methylene chloro-bromide, dichlorofluoromethane, dibromofluoromethane, bromochlorofluoromethane, dichlorodifluoromethane, dibromodifluoromethane, bromochlorodifluoromethane, trichlorefluoromethane, tribromoiluoromethane, dibromochlorofluoromethane, bromedichloroflueromethane. Two other haloethane reactants fluorinated by the present process are unsymmetrical tetrachlorcethane and hexachloroethane. Typical of the three and four carbon arom reactants which may be employed are 1.1-di-bromopropane, 1.1,1-irichloropropane, 2,2-dichieropropane, 1-bromo-1-chierobutane, 1.1,1-trichlorobutane and 2,2-di-55 chlorobutane.

Most unsaturated aliphatic chlorohydrocarbons can be reacted with gaseous chlorine and hydrogen fluoride over a basic chromium fluoride catalyst of the present invention to prepare fluorinated haloalkane reaction products containing no more hydrogen atoms than the initial chlorohydrocarbon reactant, i.e. trustion products which have not been hydrofluorinated. For example, when vapourized perchloroethylene admixed with at least 0.5

of a molecular proportion of chlorine and at least an equimolecular proportion of hydrogen fluoride is passed at a reaction temperature through a bed of a basic chromium fluoride catalyst, the same fluorochloroethane reaction products are produced as when hexachloroethane and hydrogen fluoride are reacted under similar conditions. For high conversions, perchloroethylene is usually admixed with at least an equinolecular proportion of chlorine and with about one, two, three, or four molecular proportions of hydrogen fluoride depending on whether the chief reaction product is CCI,--CCI,F, to be CCI,F-CCI,F, CCI_F-CCIF, or CCIF,-CCIF,

In some instances unsaturated aliphatic hydrocarbons can be fluorinated in accordance with the present process, this being accomplished without altering the degree of unsaturation. Suitable halohydrocarboas are 85 those having two or more halogen atoms, (viz., chlorine, bromine, or fluorine atems), at least one of which is chlorine, or bromine, attached to a carbon atom which is at least once removed from double bonded carbon atoms which are in turn attached to a total of at least one, and preferably two or more, chlorine. bromine, or fluorine atoms. Illustrative of these unsature ed aliphatic halohydrocarbons CHCI=CCICCI, $CF_1 = CC!CC!F_1$

CCL=CFCHCl₂, CF₂=CFCHB_{f2}, CHCl=CHCCl₃, CClF=CF—CCl₂—CCl₃, CCl₄F, CHCl=CF—CH₂—CCl₃ and CF₂=CCl—CCl₂—CH₃F.

Halogen atoms attached to double 100 bonded carbon atoms are not replaced by fluorine during the process of the invention unless chlorine, (or bromine), is admixed with the unsaturated halohydrocarbon, rogether with HF, as described in the preceding paragraph. 105 thereby producing saturated fluorine-containing reaction products.

The new and improved carelysts of the invention are prepared by heating a hydrated chromium fluoride to a temperature above 350°C., ordinarily in the range of from about 350° to about 750°C, in the presence of oxygen. Due to the exothermic nature of the reaction between the hydrated chromium fluoride and oxygen, however, temperatures have sometimes risen to 900°C, and even higher without deleteriously affecting the activity of the catalyst so produced. By such treatment, the hydrated chromium fluoride is at least partially converted to a basic chromium fluoride as hereinafter described. It is praential that a hydrated chromium fluoride be employed since it has been shown experimentally that the improved caralytic compositions of the invention cannot be prepared from anhydrous chromium fluoride. C-F - Liberty A. the caralytic activity of CrF cannot be no crossed by heating to a temperature of from 350° to 750°C. In the presente of avvices Ordinarily, the trihydrate of throwithm

fluoride, CrF1.3H2O, is initially employed to prepare the novel catalysts of the invention, although any of the higher hydrates may satisfactorily be used, such as the hemiheptihydrate, CrF_{1.31}/2H₂O; the tetrahydrate CrF_{1.4}H₂O; the hexahydrate, CrF, .6H2O; and the enneahydrate, CrF3.9H2O. It is the chromium fluoride trihydrate, though, which is preferred both from the standpoint of ease of preparation, i.e. availability, and case of handling. Regardless of which of the higher hydrates is initially employed, all lose water of hydration upon heating and become the trihydrate prior to or during the early stages of the activation process. The preferred chromium fluoride hydrate used to prepare the catalysts of the invention is obtained by first reacting chromium trioxide, (CrO₃), with excess strong aqueous hydrofluoric acid in the presence of 20 oxidizable organic matter, and thereafter heating the resultant reaction product to sensible dryness. A sufficient amount of oxidizable organic material should be employed in the chromium trioxide-hydrofluoric acid reaction to reduce substantially all of the chromium to the trivalent state, substances such as formalded hyde, tolume, xylene, sugar and polyethylene being satisfactory for this purpose. Ordinarily, hydrofluoric acid of about 50 to 70 per cent by weigh: strength is added to the solid chromium trioxide to slurry it, the addition of hydrofluoric acid being continued until all of the chromium trioxide is dissolved. When the dissolution is carried out in a metal container. e.g. of a magnesium metal alloy, little if any reaction occurs between the hydrofluoric acid and the chromium trioxide until the exidizable organic compound, e.g. sugar, is added. When the dissolution is conducted in a polyethylene vessel, however, enough of the polyethylene is attacked to effect the desired reaction. Upon conducting the reaction to completion and cooling the reaction product, a bright green semisolid mass is obtained which is heated to sensible dryness, e.g. at a temperature of from 90° to about 110°C. The product so dried appears by X-ray diffraction and other analyses to consist preponderantly of alpha chromic trihydrate, 2-CrF₂.3H₂O. fluoride material may, if desired, he broken or ground into fragments or granules and activated by heating in a stream of oxygen or air as hereinatter particularly described. Usually the sensibly dry hydrate is coarsely ground, e.g. to pass through a 10 mesh screen, then graphite is admixed therewith in an amount equal to about 2 net cent by weight, and the resultant mixture pelleted. It is these pellets which are then activated. Alternatively, the hydrated chromium fluoride may be slurrled with water and pasted on a carrier, such as activated charcen! or magnesium fluoride gel, and then activated, or heated to sensible dryness and ರುವ ತಮ್ಮನಡೆ.

As hereinbefore stated, the novel basic

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chromium fluoride catalysts of the invention are prepared by heating a hydrated chromium fluoride, e.g. CrF, 3H2O, in the presence of oxygen at a temperature in the range of from about 350° to 750°C. When a carbonaceous material, such as graphite, is used to prepare pellets of the hydrated chromium fluoride, it is generally desirable to carry out the activation at a sufficiently high temperature to burn off the carbon. An activation temperature in the range of from about 500° to about 600° or 650°C. is satisfactory for oxidizing carbon, although temperatures of 900°C, and higher may be employed without adversely affecting the activity of the resultant catalyst. It is advantageous to heat the chromium fluoride rapidly to the temperature at which it becomes activated, e.g. above 350°C, when highly active catalysts are desired. Prolonged preliminary hearing at lower temperatures, 85 e.g. at about 200°C, should be avoided since such treatment usually produces caralysts of lower activity containing substantial amounts of catalytically inactive Cr₂O₂.

By activation in the presence of oxygen is meant activation (1) with substantially pure oxygen gas. (2) with a gas containing molecular oxygen, or (3) with a compound which liberates or releases oxygen under activation conditions, e.g. CrO₂. Activation is usually accomplished by passing a stream of a gas comprising molecular oxygen, such as oxygen or an oxygen-containing gas, e.g. air, through a heated bed of the hydrated chromium fluoride. The initial moisture content of the 100 activating gas stream does not appear to have any effect on the activity of the resultant catalyst. Activation in a stream of oxygen. however, ordinarily produces catalysts which are catalytically active at lower temperatures, i.e. approximately 257 to 50 C. lower, than catalys's which are prepared in a stream of air. Alternatively, the hydrated chromium duotide may be activated upon first admixing it with a minor proportion of a substance, e.g. chromium trioxide, (CrO₂), which will decompose to liberate oxygen at a temperature within the range of about 350° to 750°C, and thereafter heating the mixture, preferably after pellerization, to a temperature at or above which said substance decomposes to liberate oxygen. A minture of CrF₀3H₀O and CrO₀ the latter being present in a minor proportion up to 30 per cent by weight, is desirably pelletized with graphite and then activated by heating at a temperature above 420°C, the temperature at which CrO, decomposes to liberare oxygen. No oxygen other dran thay liberated by the thermal decomposition of CrO is necessary for the activation 1949-125 Chromic oxide, viz. Cr.O., does not discongive to liberate oxygen upon heating and therefore does not aid in the activat to of hydrated chromium fluorides in accordance with the m-thod of the invention. In fact, 130

catalysts prepared by heating pellets of a mixture of CrF,.3H,O and Cr,O, at about 550° C. are catalytically less active than CtF, per se in the vapour phase reaction of CCI, and 5 HF. Furthermore, the presence of Cr₂O₃ in substantial amounts appears to affect adversely the catalytic activity of the basic chromium fluoride catalysts of the invention.

Heating in the presence of oxygen should 10 be carried out for a time sufficiently long, (at least 10 minutes), to convert at least partially some of the hydrated chromium fluoride to basic chromium fluoride, the final catalyst containing at least 0.1 per cent by weight and 15 preferably from 1 to 20 or more per cent by weight of this substance. In general, activation is substantially complete after heating at an activation temperature in the presence of oxygen for a period of 1 to 2 hours, although 20 heating may be prolonged for as long as 20 hours or more without adversely affecting

the catalytic activity.

Following activation of the catalyst by heating in the presence of oxygen, it is usually desirable, but not essential, to pass anhydrous hydrogen fluoride over the catalyst for a short time prior to using it in a fluorination reaction. This step is carried out as a precautionar; measure, i.e. to purge any residual oxygen gas 30 from the reaction vessel before introducing the halohydrocarbon, and does not appear to affect the activity of the catalyst. However, passing a halohydrocarbon over the heated catalyst in the absence of hydrogen nuoride quickly 35 reduces the activity of the catalyst. Such a catalyst of lowered activity, as well as one which has lost part of its activity through prolonged use due to carbon deposition, can easily be reactivated by burn-off in a stream of air. 40 During reactivation, as well as during the initial activation step itself, a small amount of chromic trioxide, CrO₂, is usually present in the vent gas stream.

According to the present invention there is provided a basic chronium fluoride having a composition corresponding approximately tothe empirical formula CrO,F,, said substance being a greenish tan water-insoluble solid which is amorphous to X-ray diffraction and 50 effective in catalyzing the vapour-phase reaction of hydrogen fluoride and carbon terrachloride to carbon tetrathioride at a reaction temperature below 700°C. Some crystalline hydroxy fluorities whose compositions fall 55 within the range CrF-OH), to CrF-(OH), commonly written Cr(OH,F), have at times been found to be present in small proportions.

Preparing the catalysts of the invention, as well as conducting the fluorination reaction itself, may be accomplished in a tube made of, or lined with, any suitable material such as stonel. Incomel, nickel, silver, or platinum ("Monel" and "Inornel" are Registered Trade Marks).

In carrying out the fluorination of halo-

hydrocarbens according to the present invention using the new catalyst hereinbefore characterised, the halohydrocarbon is vapourised and passed together with hydrogen fluoride through a heated bed of the catalyst at a reaction temperature in the range of from about 125 C. to 600°C. Ordinarily, however, reaction temperatures in the range of 150° to 500°C, are employed, with temperatures below 350° to 400°C, being preferred for fluorinating most of the halomethanes and the more reactive halohydrocarbons. The fluorination temperature is dependent not only upon the reactivity of the halohydrocarbon which is to be reacted with hydrogen fluoride, but also upon the products desired, the contact time, and other factors. The optimum temperature of fluorination also varies with the activity of the catalyst which in turn depends partly upon its method of preparation, and partly upon its condition 85 due to prolonged use which results in carbon surface coatings and other effects.

The ratio of hydrogen fluoride to halohydrocarbon employed in the fluorination reaction may be varied within wide limits depending on the product desired. It is sometimes advantageous to use only 0.5 mole of hydrogen fluoride per mole of halohydrocarbon. Ordinarily, however, at least one mole of hydrogen fluoride is employed per mole of the organic reactant. The preferred ratio, for making a maximum of any specific fluorinated compound, is approximately equal to, or slightly in excess of, the proportion stoichiomethally required for producing that com- 100 pound. For instance, in converting carbon tetrachloride to chlorotrifiuoremethane, at least 3 moles of hydrogen fluoride per mole of carbon tetrachloride should be employed.

Contact times of from 1 to 20 or more 105 seconds may be used in the fluorination process. although from 1 to 10 seconds are usually preferred. Contact times longer than 20 seconds are ordinately not to be desired, if for no other reason than low throughput. A cen- 110 tact time of less than I second usually results in insufficient conversion which necessitates recycling.

The fluorination reaction is generally carried out at a pressure slightly above atmospheric although both sub-atmospheric and superaemospheric pressures are operable. Aside from greater capacity per unit volume of catalyst. higher pressures are sometimes preferred to give more highly fluorinated compounds. For 120 this purpose, pressures of from 10 to 200 pounds per square inch gauge are employed.

In some instances it has been found advanthecous to carry out the vapour-phase fluoringtion in two or more stages, for example, using 125 a first stage wherein fluorination is conducted at a temperature of about 200° C, and a second stage wherein fluorination is conducted at a temperature of from 300° to 350°C. If desired, subsequent stages at higher tempera- 130

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tures may be employed.

In the case of a difficultly vapourizable halohydrocarbon reactant, such as hexaeloroethane or carbon tetrabromide, it is sometimes advanrageous to admix it with a solvent diluent, such as perchloroethylene, to aid in the vapourization, said mixture then being passed, together with hydrogen fluoride, through a heated bed of the catalyst.

The gaseous products of the reaction may be separated into their component parts by known procedures, e.g. by a series of fractional condensations and distillations, and/or water and aqueous sodium hydroxide washes and 15 drying.

After prolonged use, carbon deposits are slowly built up on the entalysts of the invention. The rate of carbon deposition is affected by several factors, such as the identity of the halohydrocarbon feed and the temperature. For example, the rate of carbon formation is more rapid in the fluorination of curbon tetrabromide than carbon tetrachloride. Furthermore, the higher the temperature at which the 25 fluorination reaction is carried out, the faster is carbon usually formed on the catalyst.

These carbon-containing catalysts of lowered activity may be regenerated as aforesaid by passing a stream of oxygen or oxygen-contain-30 ing gas over the heated carriyst at a temperature of about 500°C. Heating in the presence of oxygen is continued until carbon diexide is no longer detected in the vent gas. A stream of anhydrous hydrogen fluoride may then be passed over the catalyst to saturate the catalyst with HF prior to another fluorination run.

Following is a description by way of example of methods of carrying the invention into effect. In these examples, the terms HF effi-40 cierry, titrated conversion, and titrated organic recovery have the following meanings. The HF efficiency in mole per cent is equal to the moles of HCl in the product gases (determined by titration) times 100 divided by the meles of HF charged. The mole per cent dirated conversion of the halohydrocarbon, RXn, to the thuorohalohydrocarbon, $RX_{m-n}F_n$, is calculated as i/n times the moles of HCl. (or HBr), titrated times 100 divided by the moles of 50 halohydrocarbon charged, e.g. the titrated conversion of CCI, to CCI,F, is enfeulated as 1/2 times the moles of HCI titrated times 100 divided by the moles of CCI, charged. These terms are based on the assumption that for 55 every gram mole of HF disappearing, one gram mole of HF has been reacted with one gram mule of the halohydrocarbon reactant to form one gram mole of HCl, (or HBr). Since some halohydrocarbon reactants may be, and go frequently are, di-, tri-, or poly-fluorinated, the titrated conversion is a good indication not only of the number of gram moles of halohydrocarbon fluorinated, but also of the number of gram atoms of fluorine introduced by 65 fluorination. The weight per cent titrated

organic recovery is calculated from the total weight of the product in grams plus 16.457 grams per mole of HCl titrated, (or 60.916 grams per mole of HBr), times 100 divided by the total weight of the halohydrocarbon 70 charged.

Example 1

The vapour-phase fluorimation reaction of CCI, with HF was carried out over a pelleted basic chromium fluoride catalyst of the inven- 75 tion as hereinafter described.

The catalyst was prepared from a commercial grade high purity CrF,3H2O which was first admixed with 2 per cent by weight 80 graphite and then pressed into disc-shaped pellets 3/16 inch thick by 3/16 inch in diameter. These were loaded into a 2 inch inside diameter vertical nickel reaction tube to give a bed of CrF,3H2O pellets 24 inch deep. A 85 stream of air was then passed upwardly through the bed of pellets at a rate of from I to 2 litres per minute and the pellets were then rapidly heated to a temperature of about 500°C, in a period of about 30 minutes by 90 means of an electric furnace surrounding the reaction tube. The aforesaid flow of air and temperature were maintained constant for about 2 hours, the temperature being measured and controlled by means of a thermocouple located between the furnace and the reaction tube near the top, i.e. exit end, of the bed of pelicis.

Thereafter, the temperature was lowered, the catalyst bed fleshed with gaseous hydrogen 100 fluoride for about 30 minutes, and the following 4 runs carried out by passing a vapourphase mixture of anhydrous hydrogen fluoride and carbon tetrachloride upwardly through the catalyst bed maintained at a temperature of about 180°C. The vapour-phase reactant mixturn fed to the reaction tube was formed by bubbling anhydrous hydrogen fluoride gas upwardly through liquid carbon tetrachleride contained in a heated nickel vessel, viz. vapourizer. 110 The amount of carbon tetrachloride picked up by the hydrogen fluoride gas, i.e. the ratio of hydrogen fluoride to carbon tetrachleride, was controlled by regulating the temperature of the liquid in the vapourizer. Thus, the higher the temperature of the carbon tetrachloride, the higher was its concentration in the reactant mixture.

The effluent gas stream from the reactor was scrubbed with ice-cold water in a polysubplene vesses so as to remove the acidic constituents, viz. HCl and HF, and to condense the higher beiling constituents. Thereafter the water-scrubbed gas stream was dried by passing it first through a trap cooled in an ice-bath 125 and then through a tube of anhydrous calcium sulphate. Finally the substantially dry gas stream was passed into a refrigerated grap cooled in dry len to condense the lower boiling compenents. Unfendensed gases were collected 130

by water displacement. The scrubber water was titrated with standard sodium hydroxide and silver nitrate solutions, and from the titration values were calculated the HF efficiency, the titrated conversion, and the titrated organic recovery as previously defined. Representative samples of the organic products from all four runs were also separated into their component parts by low temperature fractional distillation io in a Podbielniak column and thereafter analyzed for product distribution. From these distillation dam, the mole per cent recovery, based on the carbon tetrachloride charged, was calculated for each component of the organic 15 product. These and other data for the 4 runs are given in Table I.

As hereinbefore stated, these runs were all carried out at a temperature of 180°C. In runs 1 to 3, the molar reactant ratios of HF 20 to CCI, were decreased from 2.3 to 1.47. In the same runs, the contact time was varied inversely as the reactant ratios. To show that reactant ratios, not contact times, were critical, a 4th run was conducted at a reactant ratio of 1.46, i.e. almost the same as that of Run 3, but at a contact time almost twice as long. Very little difference was observed upon analyzing the products of runs 3 and 4.

The catalyst employed in Runs 1 to 4 was 30 a greenish tan, water insoluble solid which was found by chemical analysis to correspond to the empirical formula CrO,F, When examined by X-ray diffraction analysis, it was found to be amorphous, i.e. no crystals could be detected by a method capable of distinguishing crystallites 100 angstroms or larger in size. No anhydrous chromium fluoride, CrF,, was observed by X-ray diffraction.

A fifth run was carried out over a CrF, catalyst not in accord with the invention prepared by passing nitrogen, not air, over a bed of high purity commercial C-F, 3H₂O pellers containing 2 per cent by weight graphite. 45 Except for employing nitrogen instead of air, the method employed in the preparation of this catalyst, the volumes, temperatures, and time were all the same as those of A above, including the final purge step with gaseous hydrogen 50 fluoride. Run number 5 was conducted according to the same general procedure and in the same equipment described in A by passing a vapour-phase mixture of HF and CCl, in a mole ratio of 1.63 over the catalyst heated to a temperature of 250°C, for a period of 30 minutes. Even at this higher temperature, the HF efficiency and the titrated conversion to CCl2F2 were considerably lower than in the preceding runs. A study of the organic recoveries after distillation shows that the product of Rim 5 actually contained less than 4 mols per cent CCLF, while greater than 50 mole per cent CCl₂F₂ was present in each of the products of the runs of A above. This and other data for Run 5 are shown in Table I.

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This catalyst was found by X-ray diffraction analysis to consist preponderantly of anhydrous chromium fluoride, CrF,.

EXAMPLE 2

The reaction of CCI, with HF was catalyzed with a fragmented basic chromium fluoride carriyst as described below.

The catalyst was prepared by first dissolving 1580 grams of CrCl, in water and admixing it with excess aqueous ammonium hydroxide to precipitate chromium hydroxide, filtering the precipitate and washing it with water, and thereafter reacting the precipiente with 1000 grams of 60 per cent by weight hydrofluoric 80

Approximately one-fifth of this green chromium fluoride reaction product was then dried to a hard cake by heating it overnight in a steam oven at 95°C. The resultant 85 CrF₁.3H₂O was broken up into fragments that passed through a 4 mesh but not through a 12 mesh screen. Some of these fragments were loaded into the same nickel reaction tube employed in Example I thereby forming a bed of CrF,3H2O fragments 2 inches in diameter by 12 inches in length. After starting a stream of moist air through the reaction tube at a rate of from 1 to 2 litres per minute, the temperature was raised to 550°C, and held there for about two hours. Therefor the flow of air was discontinued, the tem, lature of the bed lowered, and a stream of anhydrous hydrogen fluoride passed over the catalyst for a short time.

Through this bed of catalyst maintained at a temperature of 250°C, was then passed a vapour-phase mixture of HF and CCl, in a ratio of 1.41 according to the general pro-edure described in Example 1. Ail of the 105 data for this run, (Run 6), are given in Table

The catalyst employed in Run 6 was amorphous to X-ray diffraction.

Another CrF, catalyst not in accordance with the invention was prepared by passing a stream of nitrogen, not air, over some of the 4 to 12 mesh fragments of CrF, 3H2O whose preparation is described in part A above. A stream of nitrogen gas at a rate of from 1 to 2 litres per minute was passed for 4.5 hours through a 23 inch bed of these chunks heated to 450° C, in the 2 inch nickel reaction tube previously employed. After purging the cata- 120 lyst bed with anhydrous hydrogen fluoride, a vapour mixture of HF and CCI, was passed through the catalyst bed first at 550° C., (Run 7), and then at 250° C., (Run 8), the molar ratios of HF to CCI, being 1.33 and 1.68 for 125 Run 7 and Run 8 respectively.

After these runs, a stream of moist air at a rate of from 1 to 2 litres per minute was passed for two hours through the catalyst bed heared to a temperature of 550°C. Fellowing trust- 130

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ment of moist air, the temperature was lowered and the bed was flushed with gaseous hydrogen fluoride. Thereupon, a vapour mixrure of HF and CCl, in a mole ratio of 1.68 was passed for 30 minutes through the bed of catalyst at a temperature of 250°C., (Ruo 9).

The data for all four runs, (Runs 6 to 9), are given in Table I. Upon studying the data, it will be seen that the results obtained in Run 10 6, (according to the invention), are superior to those obtained in Runs 7 to 9 inclusive, (not in accordance with the invention), even though Run 7 was carried out at a temperature 300°C. higher than Run 6, (note especially the organic recoveries for these two runs). Also note that the activity of the catalyst employed in Runs 7 and 8, (prepared by heating CrF,3H,O in a stream of nitrogen gas), could not be subsequently increased by heating it in a stream of 20 air prior to Run 9.

Yet another CrF, catalyst not in accordance with the invention was prepared by first dissolving in 400 millilitres of dilute aquecus 25 hydrofluoric acid, the remaining four-fifths of the chromium fluoride reaction product of paragraph I, part A of this Example. One theusand grams of 4 to 8 mesh activated charcoal was then impregnated with this acidified 30 chromium fluoride solution. Some of the charcoal so impregnated was packed to a height of 24 inches in the nickel reaction tube and heated in a stream of nitrogen to a temperature of 450°C, at which temperature it was held 35 for about 4.5 hours. Thereafter the reaction tube was flushed with hydrogen fluoride gas and the temperature of the catalyst bed raised to 550°C.

Two runs were then carried out by passing 40 a vepour mixture of HF and CCl, over this catalyst, (not in accordance with the invention), the first run, (Run 10) being carried out at a temperature of 550°C, employing a molar reactant ratio of 1.41, (HF to CCI,), the second run, (Run 11), at 250°C, employing a reactant ratio of 1.60,

The data for Runs 10 and 11 are contained in Table I: As shown by the product data, e.g. erficiencies, conversions, and rocoveries, even 50 Run 10 which was carried out at 550°C, using a contact time of almost 18 seconds is inferior to Run 3 of Example 1, (in accordance with the invention), carried out at 180°C, using a contact time of about 3 seconds. It is also worthy of note that the activities of the catalysis of Runs 7 to 11 were of the same order of magnitude, although the surface area of the charcoal supported catalyst of Runs 10. and 11 was disproportionately greater than the fragmented catalyst of Runs 7 to 9.

> Example 3 The fluorination of CCI, with Hir was con-

ducted over a basic chromium fluoride catalyst prepared as described below.

A catalyst in accordance with the invention was prepared by reacting high purity chromium moxide, (C+O₃), with an excess of 70 per cent by weight hydrofluoric acid. The semi-crystalline bright green reaction product was heated in a drying oven at 80°C, to sensible dryness. This sensibly dry product, consisting preponderantly of =-CrF,.3H2O, was ground to pass through a 10 mesh screen, admixed with 2 per cent by weight graphite, and pressed 75 into 3/16 inch by 3/16 inch disc-shaped

These pellets were packed to a height of about 12 inches in the 2 inch nickel reaction tube employed in the previous examples. They were then activated by heating them to, and holding them for two hours at 500°C, in a stream of air. Thereafter the catalyst bed was cooled, flushed with hydrogen fluoride, and a vapour mixture of HF and CCl, in a mole ratio of 1.55 was passed therethrough at a temperature from 150° to 170°C. The data for this run, (Run 12), are contained in Table I.

Another catalyst also in accordance with the invention was prepared by passing a stream of exygen through a bed of 3/16 inch by 3/16 inch disc-shaped pellets containing 2 par cent by weight graphite prepared according to the procedure of A above. The dimensions of the catalyst bed and the conditions of the activation step were the same as above-described except that oxygen was employed instead of air. Following activation, the catalyst bed was cooled, purged with hydrogen fluoride, and a 100 vapour mixture of HF and CCl, in a mole ratio of 1.74 was passed therethrough at a temperature of about 125° to 140°C. The data for this run. (Run 13), are shown in Table I.

As will be seen by comparing the data of Runs 12 and 13, similarly good results were obtained in both runs. To produce comparable results, however, a slightly higher tentperature was required with the air-activated catalyst of Run 12 than with the exygenactivated catalyst of Run 13.

The entalysts employed in parts A and B above were amerphous to X-ray diffraction analysis.

EXAMPLE 4

The reaction of HF with CCl, was catalyzed with another basic chromium fluoride entalyst of the invention prepared from CrF,3H,O and CrO, as hereinafter described. High purity commercial CrF,3H,0, (900 grams), and CrO , '300 grams', were admixed tograther with 2 per cent by weight graphite and presed into 3/16 inch by 3/16 inch disc-shaped peliers. These pellets were packed to a height of 17

inches in the nitkel reactor employed in the previous examples and heated for 2 hours at $S(X)^{*}(\mathbb{C}_{+})$ in a stream of nitrogen gas. After cool-

ing the reactor and flushing it with hydrogen fluoride, a vapour-phase mixture of HF and CCU, in a mole ratio of 1.6 was passed through

TABLE 1

the catalyst bed at a temperature of 250°C. The reaction data for this run, (Run 14), are given in Table I.

Tri- Edification of convern. Editors Convern. Organic traited iency i.o CCL ₁ F ₃ Recovery CCLIF ₃ CCL ₃ F ₃ CCCL ₃ F ₃			Mole			Time	HC		Titrated	Titrated	Total (Organic Recovery After Bused on C.C.I. Charged	rganic Recovery Afte Bused on C.C.I, Charged		Distn.
2.3 Sec min mol %	-	115 Seed	HI. CCI.		Contact Time	of Run	Tri- trated		Convern. to CCl ₄ F ₃		CCIF,	CCI,IF3		7 100	Total
2.30 180 1.9 23 17.33 91.3 108.2 97.6 19.9 63.7 1.81 180 2.3 25 15.17 99.6 90.1 93.8 6.5 63.7 1.45 180 3.2 24 9.38 100 73.7 98.2 3.8 51.9 1.46 180 5.8 40 8.60 98.4 72.0 96.3 2.7 52.6 1.63 250 1.2 30 4.20 30.6 24.9 97.8 0.2 3.5 1.41 250 2.5 30 4.08 91.8 65.0 95.6 3.2 35.6 1.68 250 1.5 30 4.09 3.8 62.4 79.8 9.6 40.9 1.68 250 1.7 120 3.2 96.8 68.1 94.8 15.2 94.4 1.5 1.6 30 2.5 18.8 15.2 94.4	į	Zins	•	نٍ	Sec	mim	mols	% loui	mol %	wt. %	mel %	mol %	% low	% low	mol %
1.81 180 2.3 25 15.17 99.6 90.1 98.8 6.5 63.7 1.47 180 3.2 24 9.38 100 73.7 98.2 3.8 51.9 1.46 180 5.8 40 8.60 98.4 72.0 96.3 2.7 52.6 1.63 250 1.2 30 4.20 30.6 24.9 97.8 0.2 3.5 1.41 250 2.5 30 4.08 91.8 65.0 95.6 3.5 35.6 1.68 250 1.5 30 0.49 3.6 3.0 9.6 40.9 1.68 250 1.4 30 0.62 4.2 3.5 9.6 40.9 1.69 250 1.7 120 3.2 96.8 68.4 94.8 1.60 250 1.8 30 2.5 18.8 15.2 94.4 1.55 150-170 2.0 </td <td></td> <td>367</td> <td>2.30</td> <td>180</td> <td>1.9</td> <td>23</td> <td>17.33</td> <td>91.3</td> <td>108.2</td> <td>9.76</td> <td>6.91</td> <td>63.2</td> <td>10.7</td> <td>5.4</td> <td>99.3</td>		367	2.30	180	1.9	23	17.33	91.3	108.2	9.76	6.91	63.2	10.7	5.4	99.3
1.47 180 3.2 24 9.38 100 73.7 98.2 3.8 51.9 1.46 180 5.8 40 8.60 98.4 72.0 96.3 2.7 52.6 1.63 250 1.2 30 4.20 30.6 24.9 97.8 0.2 3.5 1.41 250 2.5 30 4.08 91.8 65.0 95.6 3.2 35.6 1.68 250 1.5 30 0.49 3.6 3.0 96.0 40.9 1.68 250 1.7 30 0.62 4.2 3.5 93.0 40.9 1.69 250 1.7 120 3.2 96.8 68.1 94.8 1.4 1.60 250 1.8 30 2.5 18.8 15.2 94.4 10.6 53.0 1.55 150-140 30 2.5 18.8 15.2 94.4 10.6 53.0 1.74		305	1.81	180	2.3	25	15.17	9.66	90.1	93.8	6.5	63.7	20.0	9.5	7.66
1.46 180 5.8 40 8.60 98.4 72.0 96.3 2.7 52.6 1.63 250 1.2 30 4.20 30.6 24.9 97.8 0.2 3.5 1.41 250 2.5 30 4.08 91.8 65.0 95.6 3.5 35.6 1.68 250 1.5 30 0.49 3.6 3.0 96.0 40.9 1.68 250 1.4 30 0.62 4.2 3.5 93.0 40.9 1.68 250 1.7 120 3.2 96.8 68.4 94.8 74.1 1.60 250 1.8 30 2.5 18.8 15.2 94.4 10.6 53.0 1.73 125—140 1.6 30 14.07 99.2 86.2 99.2 14.1 52.4 1.6 250 1.9 30 14.07 99.2 86.2 99.2 14.1 52.4		188	1.47	180	3.2	24	9.38	8	73.7	98.2	3.8	51.9	19.4	21.5	9.96
1.63 250 1.2 30 4.20 30.6 24.9 97.8 0.2 3.5 1.41 250 2.5 30 4.08 91.8 65.0 95.6 3.2 35.6 1.33 550 6.6 30 1.69 93.8 62.4 79.8 9.6 40.9 1.68 250 1.5 30 0.49 3.6 3.0 96.0 7.5 40.9 1.68 250 1.7 120 3.2 96.8 68.4 94.8 74.1 94.8 1.60 250 1.8 30 2.5 18.8 15.2 94.4 94.8 74.1 97.1 10.6 53.0 1.55 150-170 2.0 35 11.28 95.8 74.1 97.1 10.6 53.0 1.74 125-140 1.6 30 14.07 99.2 86.2 99.2 14.1 52.4 1.6 250 1.9 30		:75	1.46	180	5.8	40	8.60	98.4	72.0	96.3	2.7	52.6	24.1	16.5	95.9
1.41 250 2.5 30 4.08 91.8 65.0 95.6 3.2 35.6 1.33 550 6.6 30 1.69 93.8 62.4 79.8 9.6 40.9 1.68 250 1.5 30 0.49 3.6 3.5 93.0 7.7 40.9 1.41 550 17.7 120 3.2 96.8 68.4 94.8 74.1 94.8 74.1 94.8 74.1 94.8 74.1 94.8 74.1 94.8 74.1 94.8 74.1 94.8 74.1 94.8 74.1 94.8 74.1 94.8 74.1 94.8 74.1 94.8 74.1 94.1 74.1 94.1 74.1 94.1 74.1 94.1 75.4 74.1 97.1 10.6 53.0 1.74 125—140 1.6 30 9.7 75.6 60.3 99.2 14.1 52.4 1.6 250 1.9 9.7		272	1.63	250	1.2	. 30	4.20	30.6	24.9	8.76	0.2	3.5	30.4	62.3	96.4
1.33 550 6.6 30 1.69 93.8 62.4 79.8 9.6 40.9 1.68 250 1.5 30 0.49 3.6 3.5 93.0 7 1.68 250 1.4 30 0.62 4.2 3.5 93.0 7 1.41 550 17.7 120 3.2 96.8 68.4 94.8 7 1.60 250 1.8 30 2.5 18.8 15.2 94.4 10.6 53.0 1.55 150—170 2.0 35 11.28 95.8 74.1 97.1 10.6 53.0 1.74 125—140 1.6 30 14.07 99.2 86.2 99.2 14.1 52.4 1.6 250 1.9 30 9.7 75.6 60.3 98.2 14.1 52.4		68	7.	250	2.5	30	4.08	8.16	65.0	92.6	3.2	35.6	45.4	10.8	95.0
1.68 250 1.5 30 0.49 3.6 3.0 96.0 1.68 250 1.4 30 0.62 4.2 3.5 93.0 1.41 550 17.7 120 3.2 96.8 68.4 94.8 1.60 250 1.8 30 2.5 18.8 15.2 94.4 1.55 150—170 2.0 35 11.28 95.8 74.1 97.1 10.6 53.0 1.74 125—140 1.6 30 14.07 99.2 86.2 99.2 14.1 52.4 1.6 250 1.9 30 9.7 75.6 60.3 98.2		36	1.33	550	9.9	8	1.69	93.8	62.4	8.62	9.6	40.9	13.6	5.9	70.0
1.68 250 1.4 30 0.62 4.2 3.5 93.0 1.41 550 17.7 120 3.2 96.8 68.4 94.8 1.60 250 1.8 30 2.5 18.8 15.2 94.4 1.55 150—170 2.0 35 11.28 95.8 74.1 97.1 10.6 53.0 1.74 125—140 1.6 30 14.07 99.2 86.2 99.2 14.1 52.4 1.6 250 1.9 30 9.7 75.6 60.3 98.2		277	1.63	250	1.5	30	0.49	3.6	3.0	0.96					
1.41 550 17.7 120 3.2 96.8 68.4 94.8 1.60 250 1.8 30 2.5 18.8 15.2 94.4 1.55 150—170 2.0 35 11.28 95.8 74.1 97.1 10.6 53.0 1.74 125—140 1.6 30 14.07 99.2 86.2 99.2 14.1 52.4 1.6 250 1.9 30 9.7 75.6 60.3 98.2		298	1.68	250	F. 1	30	0.62	4.2	3.5	93.0					
1.60 250 1.8 30 2.5 18.8 15.2 94.4 1.55 150—170 2.0 35 11.28 95.8 74.1 97.1 10.6 53.0 1.74 125—140 1.6 30 14.07 99.2 86.2 99.2 14.1 52.4 1.6 250 1.9 30 9.7 75.6 60.3 98.2		95	1.4	550	17.7	120	3.2	96.8	68.4	94.8					
1.55 150—170 2.0 35 11.28 95.8 74.1 97.1 10.6 53.0 1.74 125—140 1.6 30 14.07 99.2 86.2 99.2 14.1 52.4 1.6 250 1.9 30 9.7 75.6 60.3 98.2		200	09.1	250	1.8	30	2.5	18.8	15.2	9.1.4					
1.74 125—140 1.6 30 14.07 99.2 86.2 99.2 14.1 52.4 1.6 250 1.9 30 9.7 75.6 60.3 98.2		2:15	1.55	150-170	ci	35	11.28	95.8	7.1.1	97.1	10.6	53.0	22.1	12.8	93.5
1.6 250 1.9 30 9.7 75.6 60.3		283	1.73	125—140	_	30	14.07	99.2	86.2	99.2	14.1	52.4	17.1	16.4	<u>i</u>
		15.7	9.1	250	6 -	30	9.7	15.6	60.3	98.2					

. Net according to the invention

Example 5

To demonstrate that substantial amounts of CF, can be produced at relatively low temperatures by reacting HF with CCI, over one of the improved catalysts of the invention, the following run was carried out in accordance with the procedure of Example 1A in the same nickel reaction tube over a 2 inch by 24 inch bed of catalyst prepared as described in the aforesaid example. The fluorination of CCI, was conducted at a temperature of 400°C. employing a molar reactant ratio of HF to CCI, of 7.74 and a contact time of 29 seconds. Throughout the run of 101 minutes, 88 grams CCI, and 88 grams of HF were fed to the reactor. The organic product was condensed and fractionally distilled. Based on the CCI, charged, a 99.7 mole per cent recovery \$1 the organic product was obtained as follows: -

...

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20	CF.	16.3 mole %
217	CCIF,	80.0 mole %
	CCI,F,	2.2 mole 😘
	CCI,F	0.7 mole %
	CCI.	u.5 mole %

EXAMPLE 6

A catalyet similar to that used in Example 3B was tested for 2200 hours in the vapourphase fluorination of CCI, with HF, mostly at a temperature of 250°C., a reactant ratio 30 of HF to CCl, of about 1.3 to 1.4, and a contact time of about 2 to 3 seconds. High HF efficiencies and organic recoveries, (95 to 100 mole per cent), were obtained throughout the run. During the run, the caralyst was burned 35 off several times with oxygen at 550°C. The catalyst showed no less in activity with burnoff, being as effective after the 2200 hour run as it was initially. No appreciable physical disintegration of the catalyst was observed.

EXAMPLE 7

40 CCl₂F and HF were reacted in the same manner and in the same nickel reaction tube over a 16 inch bed of a catalyst prepared according to the same procedure used in 45 Example 3B. During a run of 45 minutes, 1947 grams, (14.15 moles), of CCl₂F and 102 grams, (5.12 moles), of HF were passed through the cotalyst bed at a temperature of 250°C, and at a contact time of 2.9 seconds. 50 The calculated HF efficiency was 97.3 mole per cent. The organic product was separated by fractional distillation. Based on the CCIF charged, a recovery of 100 mele per cent was obtained as follows: -

55	CCIF,	1.0 mole f	·: .
	CCi ₁ F ₂	47.3 minto (. .
	CCLE	51.7 male 1	- /

EXAMPLE 8

The fluorination of CHCL, with HF was 60 carried out according to the precedure of

Example 1A in the same nickel reaction tube and over a 24 inch bed of catalyst prepared as described in the aforesaid example. The fluorination reaction was conducted at a temperature of 400°C, employing a moiar reactant ratio of 3.6 and a centact time of about 20 seconds. Throughout the run of 42 minutes, 80 grams of CHCl, and 48 grams of HF were fed to the reactor. The organic product was separated by fractional distillation. Based on the CHCI, charged, a 94.1 mole per cent recevery was obtained as follows: -

CHF,	91.0 mole 😘	
CHCIF,	3.1 mole %	
CHCL:F	0.0 mole %	75

EXAMPLE 9

The reaction of CH2Cl2 and HF was carried out in the same manner in the same nickel reaction tube and over a 17 3/4 inch bed of catalyst prepared as described in Example 3B. Through the catalyst bed heated to 250°C. were passed 372 grams CH₂Cl₂ and 243 grams HF during a period of 72 minutes. This represents a mole ratio of HF to CH₂Cl₂ of 2.8 and a contact time of 5.9 seconds. The organic reaction product was fractionally distilled. Based on the CH₂Cl₂ charged, a receivery of \$1.9 mole per cent was obtained as follows: —

CH ₂ F ₃	43.6	me!c		
CH CIF	9.1	mole	• •	90
CH ₂ Cl ₂	29.2	mole	17.5	

EXAMPLE 10

A vapour mixture of HF and CBr, in a mole ratio of 8.0 was passed at a contact time of 2.2 seconds through a 12 inch bed of the same-type catalyst and reacted according to the same procedure employed in Example 3A. During a period of 60 minutes, 865 grams of CBr, and 408 grams of HF were put through the catalyst bed at a temperature of 300°C. 100 Upon fractionally distilling the organic product, the recovery, (based on the CBr, charge). was found to be as follows: -

CBrF.	\$2.3 mole *1	
CHF,	4.5 mole *1	105
CBr ₂ F ₃	7.8 mele 1	
CDr.F	0.8 mole	

EXAMPLE II

CBr F and HF were passed as a temperature of 250°C, through a 24 inch bed of the same- 110 type chinlyst and reacted according to the same procedure employed in Example 1A. There were passed 947 grams CBr F and 150 grams of HF through the catalyst bed in approxi-mately 55 minutes. This represents a mole 115 ratio of HF to CBt F of 2.58 and a contact time of 3.1 seconds. Upon distillation of the organic product, the recovery was found to be as follows: —

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CBrF.	21.9	mole	%
CBr ₂ F ₃	54.0	mole	%
CR+ F	2.6	male	3/

Example 12

The reaction of HF and CHBr, in a mole ratio of 3.26 was carried out at 320°C. according to the same procedure and over a 24 inch bed of the same-type catalyst employed in Example 1A. During 300 minutes, 3615 grams of CHBr, and 934 grams of HF were passed through the heated bed of catalyst at a contact time of 7.8 seconds. The titrated conversion to CHBrF, was 49 mole per cent.

Example 13 CCLF—CCLF was fluorinated with HF over a 15 inch bed of the same-type catalyst and according to the same procedure employed in Example 3B. The fluorination was carried out for 60 minutes at 500°C, during which 20 time 1332 grams of CCl₂F—CCl₂F and 280 grams of HF were passed through the reaction tube. This represents a mole ratio of HF to CCl₂F—CCl₂F of 2.15 and a contact time of 2.6 seconds. The organic product was fractionally distilled and a recovery of 94.3 mole per cent, (based on the CCLF-CCLF

> CF,--CF, 1.1 mole % 4.1 mole % CF,—CCIF, 43.3 mole % CCIF,—CCIF, 29.5 mole % CCIF,—CCI,F 16.8 mole % CCL_F—CCL_F

charged), was obtained as follows: -

EXAMPLE 14

CF,-CCI,-CCI,F and HF were reacted 35 at 450°C, over a 23 inch bed of the same type cutalyst in a 3/4 inch inside diameter vertical nickel reaction tube according to the same procedure employed in Example 1A. The run was carried out for 12.75 hours at an average 40 contact time of 0.7 seconds and employing a mole ratio of HF to organic of 2.3. During the run, a total of 14,520 grams of

CF, —CCI, —CCI,F and 3240 grams of HF were passed through the reaction tube. Upon separating the organic product by fractional distillation, the total recovery, (based on the CF,-CCi2-CCI2F charged), was found to be 79 mole per cmr. The organic product was 70.9 mole per cent CF.—CCI2—CCIY2 which boiled at 72°C. at 760 mm, of mercury absolute,

What we claim is: -

1. A basic chromium fluoride having a composition corresponding approximately to the empirical formula CrO.F2, said substance being a greenish tan water-insoluble solid which is amorphous to X-ray diffraction and effective in catalyzing the uppeur-phase reaction of hydrogen fluoride and curbon terrachloride to carbon terrafluoride at a reaction temperature below 700° C.

2. A catalyst useful in promoting the

fluorination of haloalkanes by vapour-phase reaction with hydrogen fluoride, said catalyst having been prepared by contacting a hydrated 65 chromium fluoride at a temperature above 350°C, with a gas comprising molecular oxygen

3. A basic chromium fluoride catalyst consisting essentially of an amorphous powder, having been prepared by heating a hydrated chromium fluoride to a temperature in the range of 350° to 900°C, in the presence of a gas comprising molecular oxygen.

4. A method of preparing a catalyst useful in promoting the fluorination of haloalkanes by vapour-phase reaction with hydrogen fluoride, said method comprising heating a mixture of a major proportion of hydrated chromium fluoride and a minor proportion of chromium trioxide at a temperature above 420°C, for a time sufficiently long to convert at least part of the hydrated chronium fluoride to a basic chromium fluoride.

5. A method of preparing a catalyst useful in promoting the fluorination of halvalkanes by vapour-phase reaction with hydrogen fluoride, said method consisting essentially of heating a hydrated chromium fluoride to a temperature in the range of from 350° to 750°C. in the presence of oxygen.

6. A method of preparing a catalyst useful in promoting the fluorination of chlorealkanes by vapour-phase reaction with hydrogen fluoride, said method consisting essentially of hearing a bed of hydrated chromium fluoride to a temperature in the range of from 350°C. to 750°C, and passing a stream of a gas comprising a molecular oxygen therethrough for a time sufficiently long for effective amount of 100 oxygen to react therewith.

7. A method according to Claim 6 wherein the gas stream is oxygen.

8. A method according to Claim 6 wherein the gas stream is air.

9. A method according to Claim 6 wherein the hydrated chromium fluoride is CrF,.3H₂O.

10. A method according to Claim 9, wherein the CrF₃.3H₂O is the alpha hydrate.

11. A method of fluorinating haloalkanes which comprises passing a vapour-mixture of a haloalkane having from one to four corbon atoms including at least one carbon atom which is attached to a minimum of two halogens other than lodine, at least one of mid halogens being chlorine or bromine, and at least an equimolecular proportion of hydrogen fluoride at a reaction temperature in the range of from 125" to 600°C, through a bed of catalyst pre- 120 pared as in Claim 6.

 A method according to Claim 11, wherein the halcalkane is a bromerikane.

13. A method according to Claim 12, wherein the bromoalkane is earbon tetrabromide.

14. A method according to Claim 11, wherein the haloalkane is a chloroalkane.

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15. A method according to Claim 14, where in the chloroalkane is carbon tetrachloride.

16. A method according to Claim 14, where-

in the chloroalkane is chloroform.

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 A method according to Claim 14, wherein the chloroalkane is hexachloroethane.

18. A method of fluorinating aliphatic halohydrocarbons having from one to four carbon atoms including at least one carbon atom, other than a double bonded carbon atom, which is attached to a minimum of two halogens other than iodine, at least one of said halogens being chlorine or bromine, and at least 0.5 of a molecular proportion of hydrogen fluoride at a reaction temperature in the range of from 125° to 600°C, through a bed of a catalyst prepared as in Claim 6.

19. A method of preparing a fluorination

catalyst substantially as described in any one of the specific examples of the invention hereinbefore set forth.

20. A method of fluorinating aliphatic halohydrocarbons substantially as described in any one of the specific examples of the invention hereinbefore set forth.

21. A fluorination catalyst when prepared by the method claimed in any one of Claims 4 to 10 or Claim 12.

22. A fluorinated aliphatic halohydrocarbon when prepared by the method claimed in any one of Claims 11 to 18 or Claim 20.

BOULT, WADE & TENNANT, 111 & 112, Hatton Gardens, London, E.C.1, Chartered Patent Agents.

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